

Ni-functionalized Ca@Si yolk-shell nanoreactors for enhanced integrated CO₂ capture and dry reforming of methane via confined catalysis

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ABSTRACT

The emerging integrated CO₂ capture and utilization (ICCU) potentially contributes to net zero emissions with low cost and high efficiency. The catalytic performance in ICCU process is highly restricted by the equilibriums of carbonate decomposition and dry reforming of methane (DRM). Here, we engineer a unique yolk-shell dual functional nanoreactor construction to improve the catalytic performance via confined catalysis. By tailoring the carbonates decomposition kinetics and confining the CO₂ diffusion path, ~92% CO₂ conversion is achieved over (Ni/Ca)@Si and shows no distinct activity loss in 10 cycles at 650 °C. The formed Ca₂SiO₄ shells restrain the sintering of CaO yolks by acting as physical barriers, and stabilize the Ni particle size. It is also confirmed on in situ DRIFTS that the integrated DRM might occur via carbonyls, formates and CHO intermediates, in which formates species are highly dependent on Ni-carbonates interfaces.

1. Introduction

Carbon capture and utilization (CCU) technologies play pivotal roles in achieving net zero by this mid-century to relieve the climate and environmental issues caused by the greenhouse effect[1,2]. Specifically, CCU includes carbon capture (e.g., amine scrubbing[3,4] and calcium looping[5–8]) from point sources (e.g., power plants and cement factories)[9], CO₂ compression and transportation, and catalytic conversion (e.g., CO₂ methanation[10,11] and dry reforming of methane[12,13]) processes. Massive energy was consumed on temperature and pressure swing operations, severely restraining the wide deployment of such technologies[14]. Recently, an emerging integrated CO₂ capture and utilization (ICCU) technology has gained increasing interest from

researchers and engineers due to the outperformed low-cost and high efficiency[15–18]. Briefly, ICCU achieved direct conversion of captured CO₂ by gas swing between CO₂-containing exhaust gas (e.g., flue gas) and reducing agents (e.g., H₂[19,20] and CH₄[21,22]), in which the temperature-swing during adsorbents regeneration, CO₂ transportation, and CO₂ pre-heating before catalytic conversion processes are eliminated.



Among the CO₂ conversion reactions, dry reforming of methane (DRM, Eq.1) utilizing two green-house gases to yield syngas (CO+H₂), which could be further upgraded into valuable olefins and liquid chemicals via Fischer-Tropsch synthesis. Integrate CO₂ capture and

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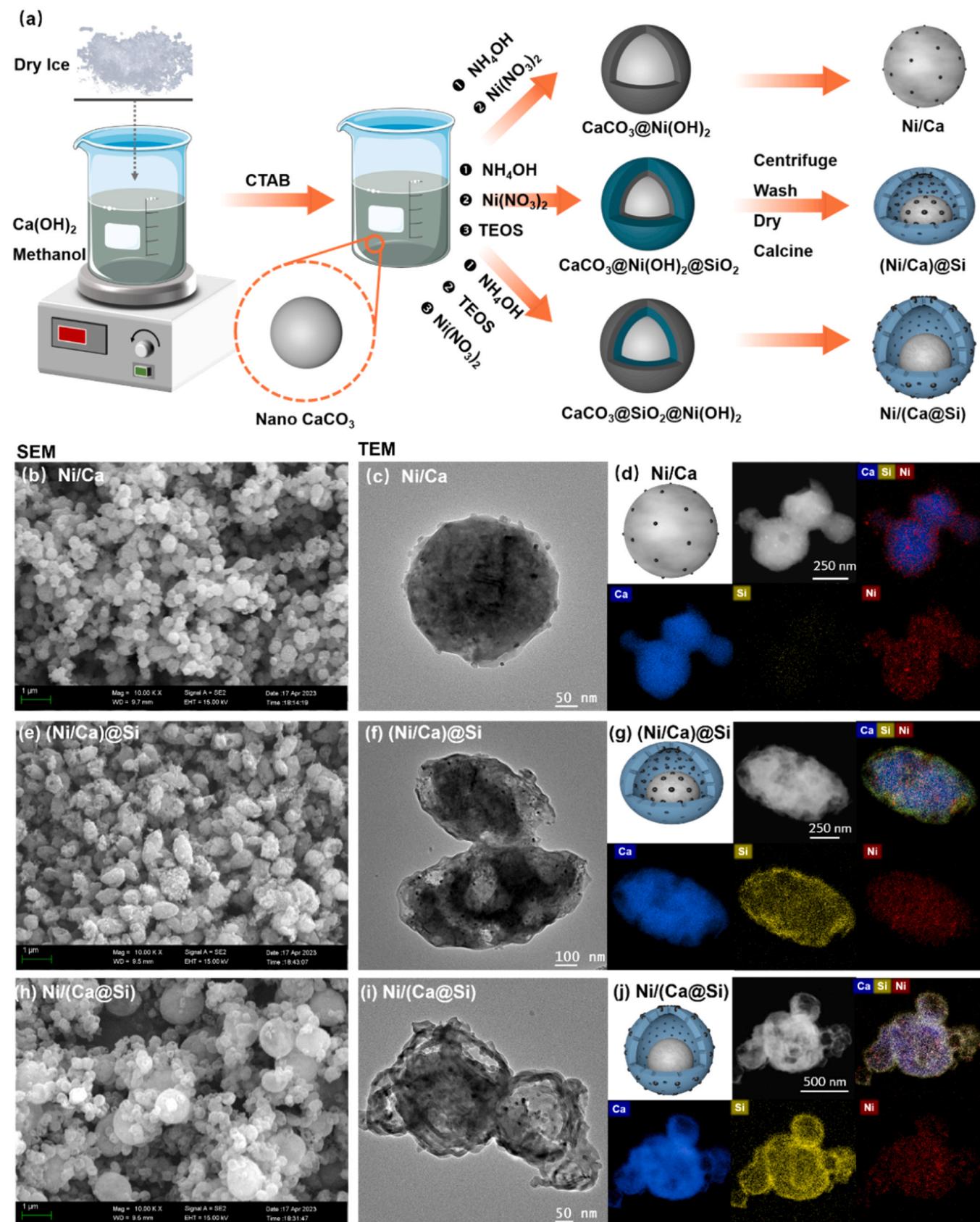


Fig. 1. Schematic diagram of the synthesis procedure of Ni/Ca and Ni functionalized Ca@Si yolk-shell nanoreactors (a); SEM and TEM-EDX-mapping images of reduced Ni/Ca (b, c and d), (Ni/Ca)@Si (e, f and g) and Ni/(Ca@Si) (h, i and j) materials.

DRM (ICCU-DRM) potentially contribute to a low-cost or profitable path for industrial CO₂ reduction[23–25]. Dual functional materials, consisting of CO₂ adsorbents and catalysts, are the keys to achieving CO₂ capture and in-situ conversion. To eliminate extra energy costs derived from temperature swings in ICCU, adsorbents and catalysts should work well under the same conditions. CaO and Ni have proven satisfactory partners for high-temperature CO₂ capture and catalytic conversion in ICCU-DRM[26,27]. However, the integrated DRM meets even more severe carbon deposition issues[24] compared to conventional DRM due to the much higher CH₄ partial pressure[28]. Furthermore, the CaO-based adsorbents exhibit significant volume swing and severe sintering during cyclic calcium looping in ICCU, which on the other hand aggravates the aggregation of loaded Ni by passive migration[21].

Abundant studies have focused on introducing thermally stable materials (e.g., ZrO₂[22] and Al₂O₃[29]) into Ni-CaO system to restrain CaO sintering for stable CO₂ capture cyclabilities, but little attention was paid on tailoring catalytic performance in the integrated DRM process. Notably, one of the reactants, CO₂, participates in DRM in ICCU via the form of carbonates, which is different from the conventional DRM process. The CO₂ exhibits a directional path (from carbonates to gas phase) in ICCU, and the decomposed CO₂ may bypass the catalytic sites and miss DRM catalytic reaction. It is hence a promising strategy to improve catalytic performance by constructing the spatial structure of the catalysts and adsorbents to effectively promote the contact probability between CO₂ and the catalyst. Confined catalysis was proposed to effectively promote the target reaction by confining reactants in an internal space[30,31] and alleviate deactivation by stabilizing the catalyst's structure[32]. The specific strategy is to build the catalytic active shell outside the carbonate core, and the decomposed CO₂ hence has to pass through the catalysts before it can be released into the gas flow. Controlled hydrolysis of silicone has reported mature strategy and makes it easy to construct porous structures[30,31]. Hence, Si was chosen as the shell material.

Furthermore, as a recently emerging process, the mechanism investigation of ICCU-DRM is far from convincing. Shao et al.[75] proposed the ¹H intermediate derived from CH₄ dehydrogenation on Ni overflow to nearby CaO surface and converted the ¹CO₂, while the ¹OH species generated from ¹CO₂ reduction participate in ¹CH₂ oxidation to form CO. Hu et al.[22] suggested that the formate species formed from adsorbed H species and CO₃²⁻ are responsible for CO formation in integrated DRM. The catalytic roles of Ni-carbonate interfaces in ICCU are also under debate[33,34]. There is a lack of a platform to systematically illustrate the independent roles of the above active candidates.

Herein, we construct Ca@Si yolk-shell adsorbents and anchor Ni in/on the shell to obtain nanoreactors. Subsequently, we demonstrate the promotion effect of CO₂ capture kinetics and stabilities and the integrated DRM catalytic performance via confined catalysis. The hypotaxis of CH₄ decomposition and integrated DRM and side reaction inhibition strategy are discussed. Furthermore, we present a convincing mechanism verification platform with various active site candidates and illustrate their independent potential routes in the integrated DRM. Both the confined catalysis-directed yolk-shell dual functional nanoreactor construction strategy and reaction pathways investigation would provide valuable guidelines to further promote ICCU-DRM performance and other heterogeneous catalytic processes.

2. Methods

2.1. Preparation of nano-CaCO₃ suspension

The nano-CaCO₃ was synthesized by dry-ice precipitation method followed previously reported work[35], as illustrated in Fig. 1a. Specifically, 1.48 g (0.02 mol) Ca(OH)₂ (Sigma-Aldrich, ≥ 95%) was dispersed in 200 mL methanol (Sigma-Aldrich, ≥ 99.5%) with stirring. Dry ice was added to the solution and the temperature was kept at ~−50 °C for 3 h. Subsequently, 30 mL cetyltrimethylammonium bromide

(CTAB, Sigma-Aldrich, ≥ 99%) solution (0.364 g in 30 mL distilled water) was added to the suspension with stirring for 10 min to avert particle aggregation. The nano-CaCO₃ was dispersed and stabilized in methanol, named as nano-CaCO₃ suspension.

2.2. Preparation of Ni/Ca, (Ni/Ca)@Si and Ni/(Ca@Si) yolk-shell dual functional nanoreactors

A certain amount (1 mL) of NH₄OH (Sigma-Aldrich, 28% NH₃ in H₂O) was added into the nano-CaCO₃ suspension with 10 min stirring to facilitate the hydrolysis of TEOS and precipitate nickel. Subsequently, 0.29 g Ni(NO₃)₂·6 H₂O (Sigma-Aldrich, ≥ 98.5%) were dissolved in 10 mL methanol and then dropped into the nano-CaCO₃ suspension and stirred for 30 min. The suspension was centrifuged and washed with methanol three times and then dried at 70 °C for 2 h under vacuum. Finally, the sample was calcined at 800 °C for 1 h with a heating rate of 2 °C/min to obtain Ni/Ca.

The (Ni/Ca)@Si and Ni/(Ca@Si) yolk-shell dual functional nanoreactors were synthesized by introducing an extra silica coating process (Fig. 1a). Briefly, 1 mL NH₄OH (a.q.) was dropped into the nano-CaCO₃ suspension with 10 min stirring. Then, 0.29 g Ni(NO₃)₂·6 H₂O (dissolved in 10 mL methanol) was added into the nano-CaCO₃ suspension and stirred for 10 mins, followed by dropping 10 mL 10 mol% tetraethyl orthosilicate (TEOS, Sigma-Aldrich, ≥ 98%) into methanol solution with 3 h stirring. The suspension was then washed, centrifuged, dried and calcined as abovementioned processes to obtain calcined (Ni/Ca)@Si. The calcined Ni/(Ca@Si) was prepared by swapping the adding order of Ni(NO₃)₂·6 H₂O and TEOS. Before the evaluation, the NiSiCa-based materials were reduced at 700 °C for 1 h in 5% H₂/Ar to obtain reduced materials.

2.3. Materials characterizations

Inductively coupled plasma-optical emission spectrometry (ICP-OES): The materials were digested in nitric acid under microwave and the Ni, Si and Ca contents were analyzed by ICP-OES (Agilent ICPOES730).

N₂ physisorption: The porosities of materials were characterized by isothermal N₂ adsorption-desorption at 77 K on Autosorb iQ Station. And the micro and meso pore size distributions were analyzed by Horváth-Kawazoe (HK) and Barrett-Joyner-Halenda (BJH) methods using the desorption branch, respectively.

Electron microscopy: The surface morphologies of the synthesized nanoreactors were monitored on a scanning electron microscope (ZEISS Merlin® FE-SEM). The detailed morphologies and element distribution were characterized by transmission electron microscope on JEOL JEM-2010 with energy dispersive spectroscopy (EDS). The Ni particle size distributions were measured and plotted by Nano Measurer 1.2. The average particle diameter (d_{ave}) was calculated based on Eq.2.

$$d_{ave} = \frac{\sum(n_i d_i^3)}{\sum(n_i d_i^2)} \quad (2)$$

X-ray diffraction (XRD): The phase composition was characterized by XRD using a PANalytical Empyrean X-ray Powder Diffractometer (45 kV and 40 mA), equipped with a Cu K α x-ray source. The data was collected in the 10–90° 2 Theta range and analyzed on MDI Jade 6.0.

Temperature-programmed CO₂ adsorption-desorption: The isothermal and programmed thermogravimetric analysis (TGA) were collected on a Hi-Res TGA 2950 thermogravimetric analyzer. Specifically, the temperature-programmed CO₂ adsorption-desorption analysis was carried out in 100 mL/min 15% CO₂/N₂ from 100 to 850 °C with a heating rate of 10 °C/min. The isothermal TG experiment was performed at 650 °C in 100 mL/min 15% CO₂/N₂ for 25 min, followed by 100 mL/min N₂ (BOC.co, 99.9%) for 45 min. To exclude the change of CaO-based materials under atmosphere storage (steam or CO₂

adsorption), all the materials were *in situ* pre-calcined in N₂ at 800 °C for 30 min.

X-ray photoelectron spectroscopy (XPS): The surface/interface element compositions and states of the synthesized nanoreactors were determined by XPS on Thermo Fisher ESCALAB 250Xi with Mg K α X-ray source.

Raman spectroscopy: The deposited carbon species were presented by Raman spectroscopy on the WITec Alpha 300 R Confocal Raman Microscope equipped with a 532 nm diode laser (10 mW). The spectrums of each material was generated from three different sample areas.

2.4. ICCU-DRM evaluation

The ICCU-DRM reactions with nanoreactors were evaluated in a fixed-bed quartz wall reactor (Figure S1, OD: 12 mm; ID: 10 mm; L: 600 mm). 0.5 g material was fixed in the middle of the reactor by quartz wool and heated by a tubular furnace (Elite TSH 12/50/300–2416CG). Moreover, the thermocouple was placed in the middle of the reactor to control the temperature.

The ICCU-DRM experiment was performed by gas swing among 15% CO₂ (CO₂ capture step), N₂ (purge step) and 10% CH₄ (DRM step). Specifically, the material was heated to 700 °C and *in situ* reduced under 100 mL/min 5% H₂/Ar for 1 h (pre-processing). The ICCU mainly includes two steps for CO₂ capture and *in situ* DRM. In the first step, 100 mL/min 15% CO₂/N₂ was introduced for 20 min, followed by N₂ purge for 3 min. In the second step, 10% CH₄/N₂ was switched into the reactor until no more CO was generated. The cyclic ICCU-DRM evaluations were exhibited by repeating the above steps with an extra 5 mins' N₂ purge among each cycle. The outlet gas was monitored by an online gas analyzer (Enerac 700 AV) equipped with a Non-Dispersive InfraRed (NDIR) detector and H₂ thermal conductivity detector (CIXI-CX-H02).

The quantitative analysis was calculated by integrating the signal of reactants and products in different step regions. Notably, the carbon deposition in the DRM step would retain to the following CO₂ capture step and then compete with CO₂ capture via the reverse Boudouard reaction (Eq. 3). The CO₂ capture capacities (C_{CO2}) were hence calculated by subtracting half of CO production from CO₂ consumption in CO₂ capture step (Eq. 4). (CO^b: benchmark signal)



$$C_{CO2}(mmol/g) = \frac{\int (CO_2^b - CO_2 - 0.5 * CO) dt * 1.667mL/s}{22.4mmol/mL * 0.5g} \quad (4)$$

The CO (Y_{CO}) and H₂ (Y_{H2}) yields were integrated in the DRM step as follows:

$$Y_{CO}(mmol/g) = \frac{\int CO dt * 1.667mL/s}{22.4mmol/mL * 0.5g} \quad (5)$$

$$Y_{H2}(mmol/g) = \frac{\int H_2 dt * 1.667mL/s}{22.4mmol/mL * 0.5g} \quad (6)$$

Due to the detection limitation on steam, the CO₂ conversion rate (X_{CO2}) was calculated based on CO₂ capacities and unreacted CO₂ (Y_{CO2}) in the DRM step, as listed below:

$$Y_{CO2}(mmol/g) = \frac{\int CO_2 dt * 1.667mL/s}{22.4mmol/mL * 0.5g} \quad (7)$$

$$X_{CO2}(\%) = \frac{C_{CO2} - Y_{CO2}}{C_{CO2}} * \% \quad (8)$$

The CO₂ removal efficiency represents the extent to which CO₂ is removed. It was calculated via below equation:

$$R_{CO2} = \frac{CO_2^{input} - CO_2^{output}}{CO_2^{input}} * 100\% \quad (9)$$

2.5. *In situ* diffused reflectance infrared fourier transform spectroscopy (DRIFTS)

The *in situ* DRIFTS was carried out on PerkinElmer Spectrum equipped with a Mercury Cadmium telluride (MCT) detector (4 cm⁻¹ resolution, 1000–4000 cm⁻¹ range). To identify the surface intermediates in the ICCU-DRM process, two sets of *in situ* DRIFTS experiments were exhibited: isothermal ICCU-DRM and temperature-programmed DRM.

2.5.1. Isothermal ICCU-DRM *in situ* DRIFTS

The materials were *in situ* reduced in the cell under 5% H₂/Ar at 700 °C for 30 min and then equilibrated to 650 °C in Ar for 10 min. The spectrum was collected as the benchmark and subtracted, and subsequently, the 15% CO₂/N₂ was introduced for 90 s as the CO₂ capture step. And then switch into N₂ for 300 s to purge the residual CO₂ and unstable surface species, followed by introducing 10% CH₄/N₂ for 500 s in the DRM step. The spectra were collected every 10 s from inletting CO₂ to exhibit the surface species changes under ICCU-DRM conditions.

2.5.2. Temperature-programmed DRM *in situ* DRIFTS

The integrated DRM was investigated under temperature-programmed conditions (100–750 °C). The materials were *in situ* reduced as above, followed by carbonating at 650 °C for 30 min under 15% CO₂/N₂. The cell was cooled down to 100 °C in 15% CO₂/N₂ and then purged for 10 min under N₂ for benchmark subtraction. Subsequently, 10% CH₄/N₂ was introduced with a heating rate of 50 °C/3 min up to 750 °C.

3. Results and discussion

3.1. The construction of Ni-functionalized Ca@Si yolk-shell nanoreactors

The CaCO₃ easily re-crystallizes in synthesis and yields large and anisotropic crystals[36], challenging the coating process in core-shell structure construction. In order to synthesize CaCO₃ with uniformly spherical morphology, introducing continuous disturbance[37] during the carbonates precipitation process to build miscellaneous crystals and lowering the temperature[38] to suppress recrystallization are considered to be effective strategies. Herein, dry ice was applied as the CO₂ source and provided satisfactory disturbance in the sublimation process in carbonate synthesis (Fig. 1a). As shown in Fig. 1b and c, the above-mentioned synthesis strategy successfully directs the construction of spherical particles, and the Ni could be uniformly deposited in such a one-pot synthesis procedure. Further construction of core-shell nanoreactors inspired by confined catalysis requires the hydrolysis of silane on the surface of calcium carbonate (Fig. 1a). CTAB was introduced after carbonate precipitation to stabilize suspension[39] and subsequently acted as the template in silica deposition[40]. As shown in Fig. 1e-j, although silane would inevitably form micelles spontaneously to form SiO₂ cavities, the silica coating on calcium was well constructed. Furthermore, the volume shrinkage of calcium carbonate decomposition [41] and the formation of Ca₂SiO₄ shell during calcination *in situ* induced the formation of yolk-shell structures.

Notably, the various coating sequences of nickel and silica derived different nanoreactor morphologies. Loading silica after nickel induces spindle-shaped nanoreactors (Fig. 1e and f), while reverse deposition order constructs spherical structures with larger cavities (Fig. 1h and i). The induction mechanism is unclear, which might be attributed to the carbonate surface modification by nickel deposition or the interaction of calcium and silica in the calcination process. However, in any case, the successful encapsulation of calcium by silicon provides a convincing platform for subsequent performance and mechanism investigation. And the presented synthesis approach allows a wide modification for producing materials with various characteristics, such as adjusting silane

Table 1

Elemental analysis and porosity characterizations of NiSiCa-based materials.

Materials	ICP (wt%)			S _{BET} (m ² /g)	V _{micro} (m ³ /g)	V _{meso} (m ³ /g)
	Ni	Si	Ca			
Ni/Ca	2.78	—	34.13	12.56	0.0031	0.088
(Ni/Ca) @Si	2.26	4.47	38.00	28.72	0.0097	0.35
Ni/ (Ca@Si)	2.28	3.47	31.01	23.57	0.0082	0.13

amount for different shell thickness.

3.2. Characterizations of Ni-functionalized Ca@Si yolk-shell nanoreactors

Apart from the morphology analysis, more detailed characterizations were carried out to fully describe the synthesized nanoreactors. As evidenced by ICP results in **Table 1**, the three materials were synthesized with identical Ni loadings to exclude the performance differences caused by various active phase content. The Si and Ca fractions were also well controlled over two different nanoreactors. XPS was applied to evaluate the surface element fraction (~1–10 nm depth) [42], as detailed in **Figure S2** and **Table S1**. The silica of (Ni/Ca)@Si and Ni/(Ca@Si) concentrated on the surface, as observed from the TEM results (**Fig. 1g** and **j**). Notably, the Ni contents on these two nanoreactors were similar (1–1.1 at%, **Table S1**), indicating the Ni distributed closely to the thin shell (**Figs. 1f** and **i**).

The phase fractions of various materials were analyzed by XRD, as shown in **Figs. 2a** and **b**. After calcination of as-synthesized materials, most of the carbonates decomposed and part of CaO reacted with SiO₂ to form the Ca₂SiO₄ phase. Notably, as evidenced in the internal pattern in

Fig. 2b, the Ni/(Ca@Si) exhibited higher Ca₂SiO₄ crystallinity compared to (Ni/Ca)@Si. It is speculated that loaded Ni would physically obstruct the direct contact of calcium and silica, hence restricting the formation of huge Ca₂SiO₄ crystalite phase on (Ni/Ca)@Si. The binding energy shift in XPS also verified the formation of Ca₂SiO₄. The electron density and electronegativity of atoms exhibit various shielding effects of core electrons[43]. Introducing atoms with higher electronegativities (e.g. X_{Si}=1.7) would attract electrons from bulk atoms (e.g. X_{Ca}=1) and weaken the shielding effect on core electrons, hence increasing the binding energy of bulk elements. The Ca 2p XPS (**Figure S2e**) peaks of (Ni/Ca)@Si and Ni/(Ca@Si) shift to higher binding energy by ~0.6 eV, indicating the extensive combination of silica and calcium (Si-O-Ca in the form of Ca₂SiO₄) on the surface. More clearly, the O1s (**Figure S2f**) of nanoreactors splits into two peaks at 530.5 and 531.5 eV, assigning to Ca-O-Ca and Ca-O-Si, respectively. The NiO was detected as the only Ni-containing crystal phase (**Fig. 2a**), consisting of Ni 2p characterization in XPS (**Figure S2c**, no distinct shift excluding the possibility of the widespread existence of Ni-O-Si).

The residual CaCO₃ on NiSiCa-based materials were thoroughly consumed in 5% H₂/Ar reduction at 700 °C, evidenced by the disappearance of CaCO₃ peaks in **Fig. 2b**. However, the NiO reduction is not obvious, which might be attributed to the encapsulated NiO species with poor H₂ accessibility. Furthermore, a small amount of Ca(OH)₂ was formed in the material due to air exposure during materials storage.

The coated silica acts as the shell of the nanoreactors, which needs to achieve both good confinement and mass transfer of reactants and products. The porosity of the shell is hence pivotal for such unique requirements. As listed in **Table 1**, the synthesized Ni/Ca as a benchmark exhibits poor porosity with 12.56 m²/g surface area, 0.0031 m³/g and 0.088 m³/g micro/mesopores volumes, respectively. The uniform CaO spheres exhibit ~300 nm diameter, which could provide ~5.97 m²/g

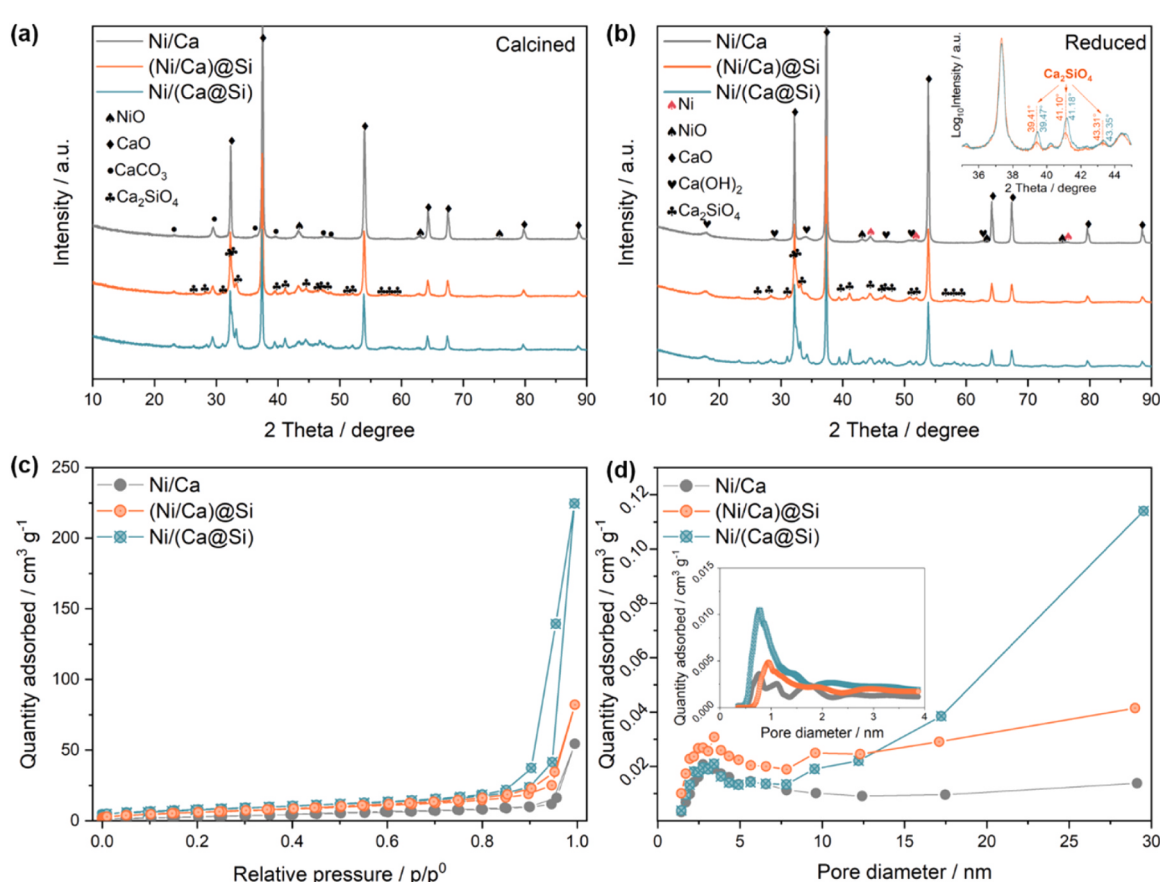


Fig. 2. The XRD patterns of calcined (a) and reduced (b) materials; The N₂ adsorption–desorption isotherms (c) and pore size distributions (d) of calcined materials.

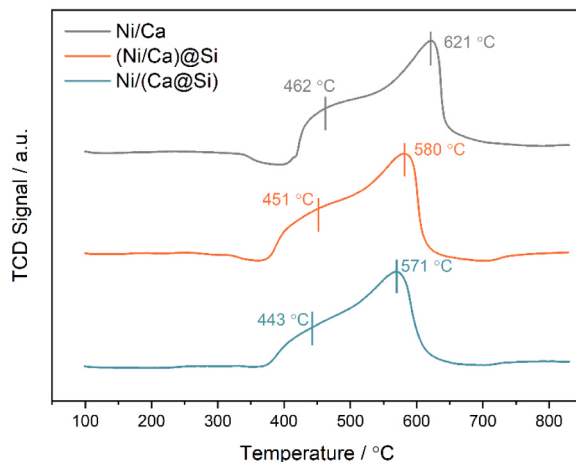


Fig. 3. TPR curves of calcined Ni/Ca, (Ni/Ca)@Si and Ni/(Ca@Si) materials.

surfaces (CaO density: 3.35 g/cm³). Apparently, the tested surface area is higher than the theoretical value, which is explained by the defects and cracks observed in SEM images (Fig. 1b). As a comparison, the (Ni/Ca)@Si and Ni/(Ca@Si) nanoreactors perform doubled surface area and multiple micro/mesopores volumes, indicating the formation of porous shells. As shown in Figs. 2c and d, the nanoreactors exhibited distinct hysteresis loops in adsorption-desorption isotherms and enhanced micro/mesopores distributions. Specifically, the Ni/(Ca@Si) shows more pores with the diameter of 10–30 nm (Fig. 2c), which is highly attributed to the abundant cavities as discussed in Section 3.1.

The reducibility of metal oxides indicates the existing reducible species, which is an important indicator of metal-support interaction

and metal distributions. As shown in Fig. 3, the H₂-TPR further confirms the existence of similar Ni species based on analogous H₂ consumption curves. Specifically, Ni/Ca exhibits two main reduction peaks at 462 and 621 °C, which are respectively attributed to the reduction of free and metal-support interacted NiO species[44,45]. It is controversial about the existing form of interacted NiO carried on irreducible supports. Some researchers suggest that the NiO reacts with MO_x to form Ni_xMyO_z spinel to present strongly interacted NiO[46,47], while others deem that the interaction could be defined by encapsulation or even contact in the absence of spinel formation[48,49]. Based on the previous XRD and XPS analysis, the nickel presents in the NiO form on the synthesized nanoreactors and no Ni-containing spinel (i.e. NiAl₂O₄) formation was identified. It is hence reasonable to conclude that the NiO reduction peak at higher temperatures is contributed from the encapsulated NiO by CaO or Ca₂SiO₄. The metal particle size is widely believed inversely proportional to the reduction temperature[50,51]. However, as summarized in Figs. S3a-c, the average Ni particle sizes are positively related to the reduction temperature of both reduction peaks. It is inferred that the degree of encapsulation on NiO species might decrease from Ni/Ca to NiSiCa-based nanoreactors, indicating that the CaO exhibits more severe encapsulation effect on NiO.

3.3. ICCU-DRM performance

The ICCU has two steps: (i) CO₂ capture and (ii) in situ catalytic conversion. The temperature is highly sensitive for CO₂ capture using CaO-based adsorbents[52,53]. As shown in Fig. 4a and b, the materials performed differentiated carbonation kinetics. Ni/Ca started to capture CO₂ from ~ 500 °C, while (Ni/Ca)@Si and Ni/(Ca@Si) nanoreactors could carbonate at even ~ 400 °C. It was reported that smaller CaO particles had promoted carbonation kinetics at low temperatures due to

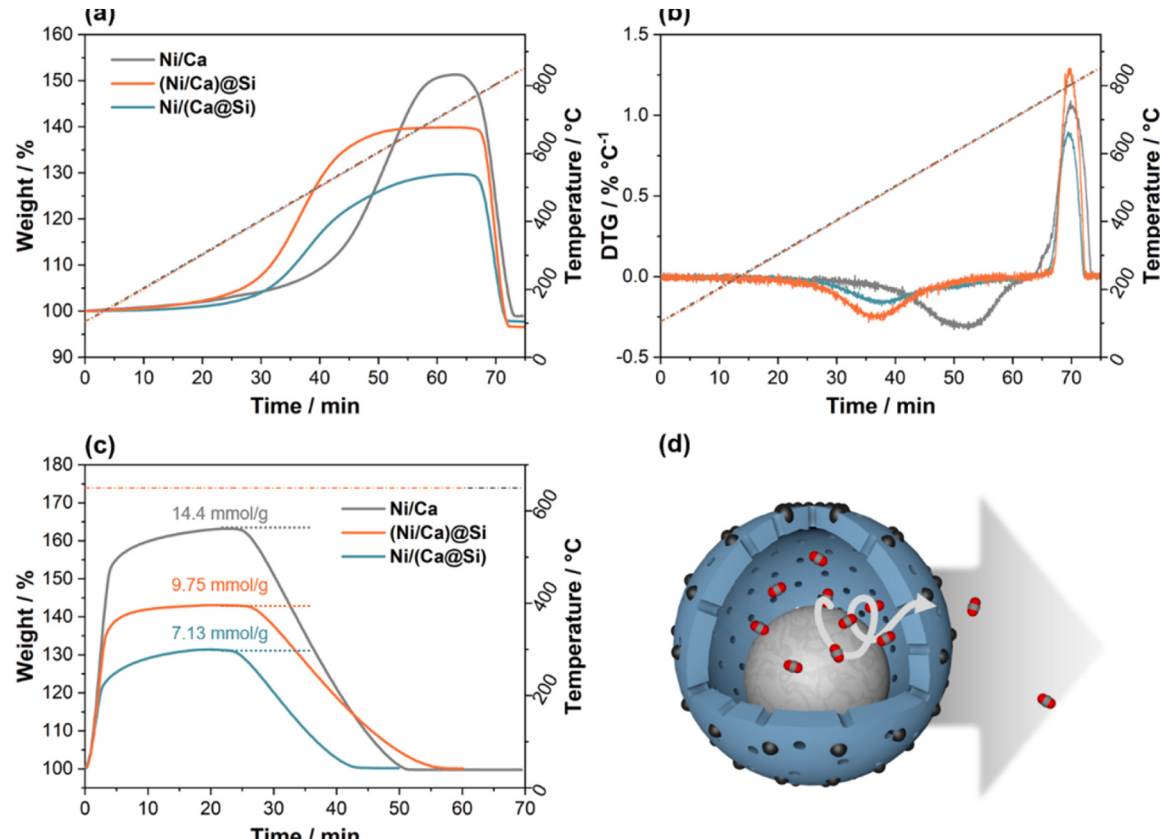


Fig. 4. The temperature programmed CO₂ adsorption-desorption curves of calcined materials: (a) TGA and (b) DTG; (c) The isothermal CO₂ adsorption-desorption curves of calcined materials; (d) The schematic diagram of CO₂ desorption on Ni/(Ca@Si) nanoreactor.

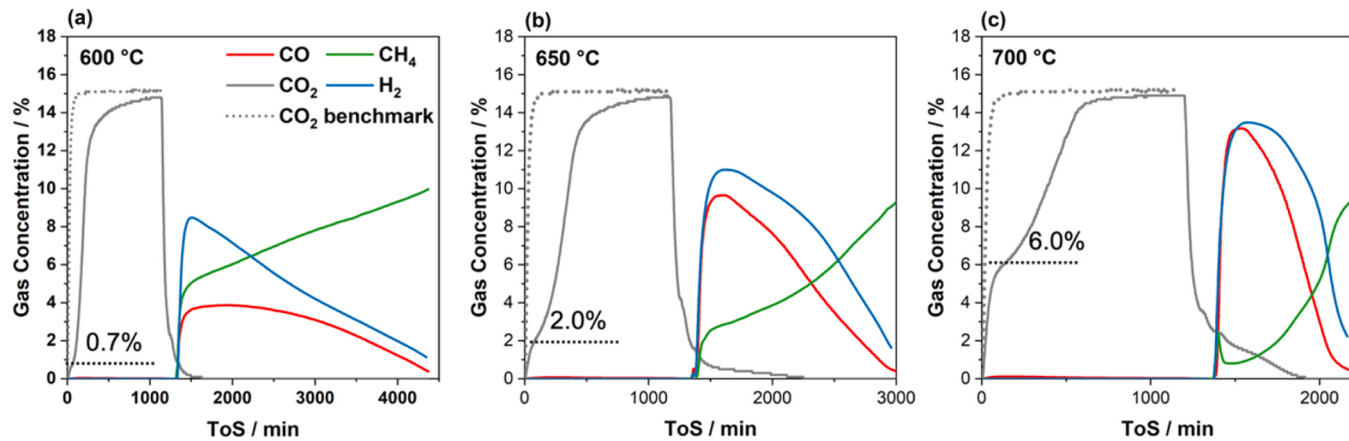


Fig. 5. Real time ICCU-DRM performances under various temperatures using Ni/(Ca@Si) nanoreactors: (a) 600 °C; (b) 650 °C and (c) 700 °C.

a more abundant surface and shorter diffusion distance[54]. As detailed in Fig. 1, the significantly smaller and looser CaO core after silica coating (Figs. 1f and i) fundamentally contributes to such enhancement in CO₂ capture. With the temperature increasing, the formed carbonates started to decompose quickly from ~750 °C, and all three candidates performed similarly. A proper temperature of ICCU should permit satisfactory carbon capture efficiency in the CO₂ capture step and appropriate decomposition kinetics in the integrated DRM step. Operating ICCU at a lower temperature would restrict the overall process efficiency, while

applying too high temperature would excessively accelerate decomposition and limit the CO₂ conversion rate. As evidenced in Fig. 4, in which the carbonation kinetics reached the peak at ~500 °C and decomposition significantly occurred from ~700 °C, it is hence recommended that 500–700 °C is suitable for CO₂ capture using CaO-based materials.

Besides the enhanced CO₂ capture performance at lower temperatures, the (Ni/Ca)@Si and Ni/(Ca@Si) nanoreactors exhibited the expected decomposition results under 650 °C isothermal conditions. As shown in Fig. 4c, the Ni/Ca exhibited the highest CO₂ capacity (~

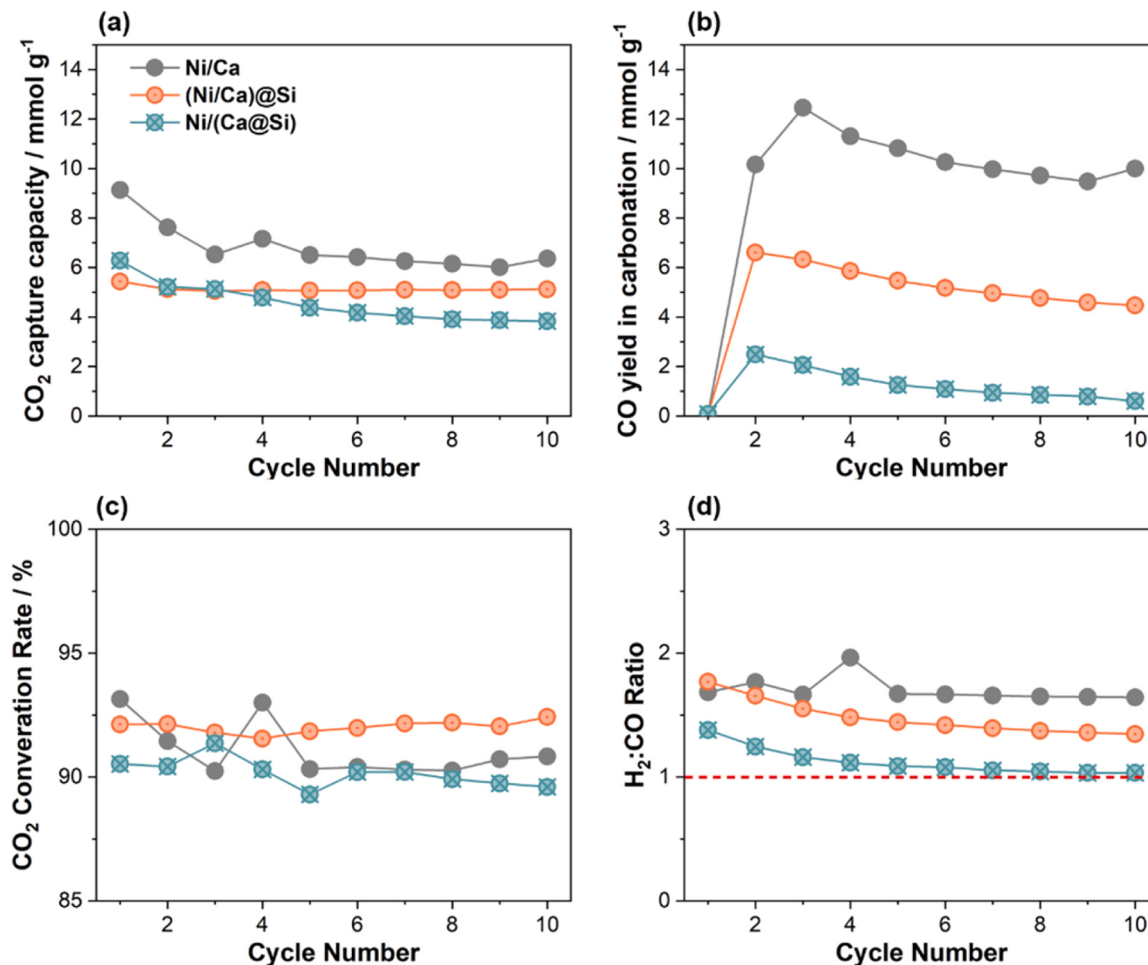


Fig. 6. Cyclic ICCU-DRM performances of the materials in 10 cycles: (a) CO₂ capture capacities and (b) CO yield in the CO₂ capture step; (c) CO₂ conversion and (d) H₂:CO ratio in the DRM step.

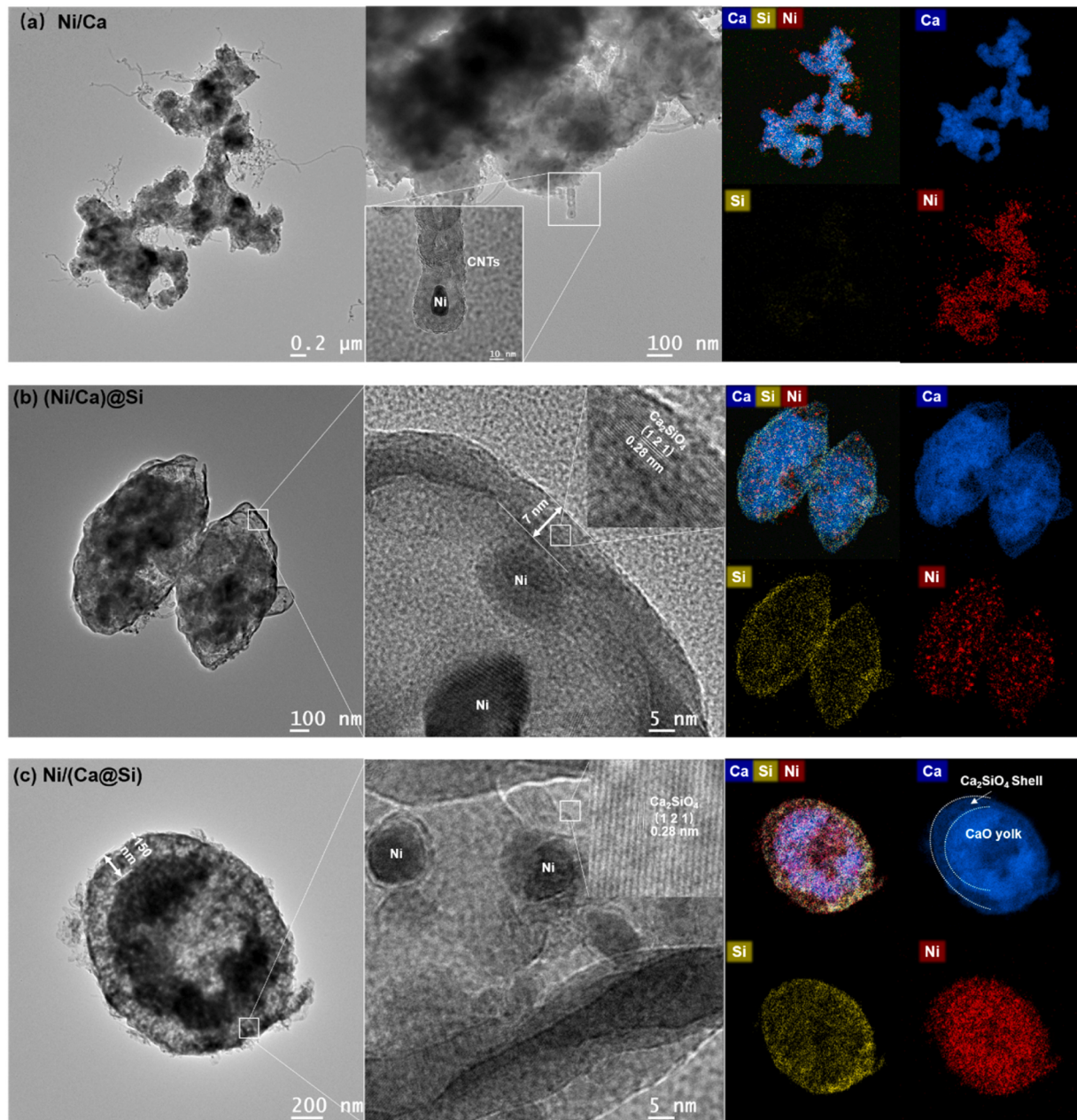


Fig. 7. TEM-EDX-mapping images of spent (a) Ni/Ca, (b) (Ni/Ca)@Si and (c) Ni/(Ca@Si) nanoreactors after 10 cycles.

14.4 mmol/g) due to the highest CaO molar fraction. Introducing silica would decrease CaO fraction and consume CaO by forming Ca_2SiO_4 , hence showing lower CO_2 capacities (~ 9.75 and 7.13 mmol/g for (Ni/Ca)@Si and Ni/(Ca@Si), respectively). Notably, the carbonates decomposition of nanoreactors is slower than the Ni/Ca benchmark, which indicates the successful preparation of confined nanoreactors. As illustrated in Fig. 4d, the porous shell provided confined cavities and extended the residual time of CO_2 after decomposing from carbonates. Riveting the catalyst on the shell would reasonably increase the contact probability between CO_2 and the catalyst, promoting the catalytic performance.

The temperature effect is further evaluated under ICCU conditions to obtain the real-time CO_2 capture kinetics and the following DRM catalytic performance. The CO_2 removal efficiency represents the extent to which CO_2 is removed in the flue gas. As shown in Fig. 5, the initial CO_2 removal efficiency of Ni/(Ca@Si) decreases from ~ 95% at 600 °C to ~ 60% at 700 °C. The equilibrium of Eq.8 shifts to carbonate

decomposition under higher temperatures due to the endothermic properties of the decomposition process, which was also evidenced by the increased CO_2 release at higher temperatures in the DRM step. The DRM, as a strongly endothermic reaction, favors high temperatures [12]. Notably, the CO_2 is restricted by carbonate decomposition, hence interfering with the effect of temperature on the integrated DRM. However, although a low process temperature ensures better CO_2 removal in the CO_2 capture step, ~ 3000 s is needed to thoroughly consume the carbonates in the DRM step. Overall, proper process temperature should fit satisfactory CO_2 capture performance and synergy carbonates decomposition and DRM kinetics. Considering the further deployment of ICCU, the residence time of CO_2 capture and DRM steps should be equal for continuous operation under mature dual fluid bed systems. Hence, 650 °C was selected as the optimal ICCU temperature with almost equal time periods of CO_2 capture and DRM.



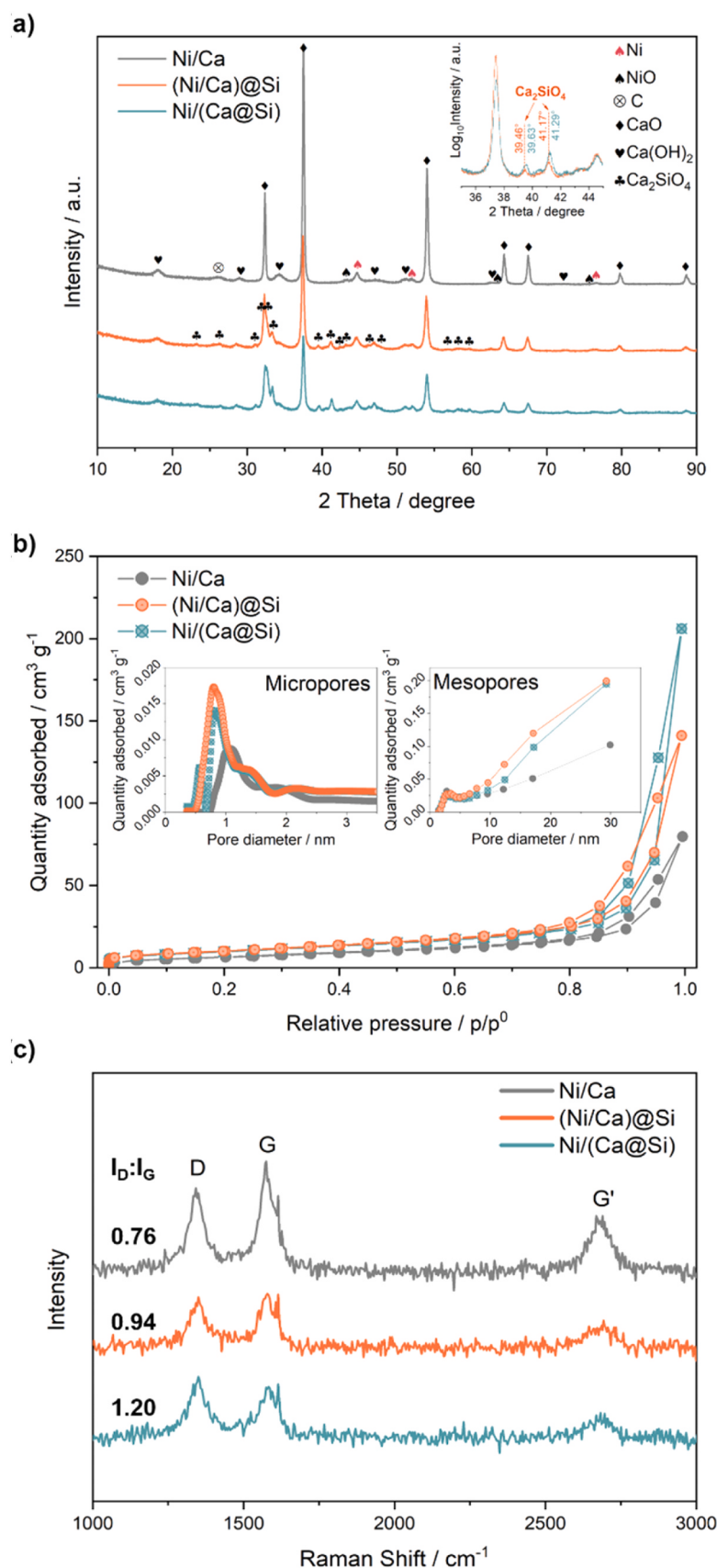


Fig. 8. (a) XRD patterns, (b) N₂ adsorption–desorption isotherms with micro-meso pore size distributions and (c) Raman spectrums of spent materials.

Ten cyclic ICCU evaluation was carried out to evaluate the stabilities of Ni/Ca, (Ni/Ca)@Si and Ni/(Ca@Si) (Fig. 6 and S4). As a benchmark, the CaO without Ni functionalization exhibits no activity in ICCU-DRM (Figure S5). In the integrated DRM process, the CH₄ decomposition inevitably occurs as a side reaction, and deposits carbon on the materials [21,24] (Fig. 7a as an example). The carbon would be then gasified by CO₂ in the carbon capture step (Figure S4) and obstruct the quantification of CO₂ capacities. Notably, the reverse Boudouard reaction might compete with carbonation in the initial time of CO₂ capture; however, 20 mins of carbonation is enough for saturating the adsorbents. The integrated CO₂ capacities in Fig. 6a were calculated by deducting half of CO production from CO₂ consumption, as detailed in Eq.4. The (Ni/Ca)@Si showed stable CO₂ uptake capacities, only decreasing from 5.43 to 5.12 mmol/g in cyclic evaluations (Fig. 6a), and the nanoparticles remain separated (Fig. 7b). The thickness of the Ca₂SiO₄ shell on (Ni/Ca)@Si was well maintained (Figs. 1g and 7b), implying the nickel intermediate layer does well in controlling the contact of calcium and silica. As a comparison, the Ni/Ca exhibited the highest initial CO₂ capacity due to the highest CaO fraction and gradually decreased from 9.13 to 6.36 mmol/g in 10 cycles due to sintering (significant particle aggregation in Fig. 7a). Similarly, the Ni/(Ca@Si) performed a capacity loss from 6.28 to 3.83 mmol/g. As shown in Fig. 7c, although the nanoparticles the Ca₂SiO₄ shell on Ni/(Ca@Si) significantly thickens to ~150 nm, which might hinder CO₂ diffusion and capacities. The increase of the thickness might be attributed to the shell reconstruction in cyclic evaluation, especially the merge of small Ca₂SiO₄ species (Fig. 1h)

outside the Ni/(Ca@Si) nanoreactors. As a comparison, the surface of (Ni/Ca)@Si is cleaner, which might contribute to the stable shell in cycles. It is reasonable to infer that the well-designed shell construction is an effective strategy to defer the CaO sintering and hence promote the CO₂ capture stabilities. Notably, the carbon deposition would harm the following CO₂ capture step by diluting the toxic CO in the exhaust gas from a process point of view. Introducing an extra intermediate step, e.g., steam purge[55], might be applicable to eliminate the CO emission and by-producing syn gas in the deployment of ICCU-DRM.



The metal particle size of Ni was believed to be sensitive in CH₄ decomposition (Eq.9) and carbon nanotube growth[56,57]. Larger Ni clusters are more prone to generate carbon deposition in DRM[58]. Hence, even though the Ni on Ni/(Ca@Si) might be more easily exposed in CH₄, it still exhibits less carbon deposition due to the best dispersion. As summarized in Figure S3, Ni/Ca exhibited the largest initial particle sizes (~ 13.31 nm) and performed the highest CO yield in the CO₂ capture step in the 2nd ICCU cycle (Fig. 6b). With the decreasing of Ni sizes from 12.12 nm on (Ni/Ca)@Si to 9.96 nm on Ni/(Ca@Si), the CO yield in the CO₂ capture step (twice the amount of carbon deposits) reduced from 6.61 to 2.49 mmol/g. Interestingly, the carbon deposit amounts decrease with cycles, while the statistical metal particle sizes increase to varying degrees. The H₂:CO exhibited a similar decreasing trend, representing the CH₄ decomposition was gradually hindered in cycles. Notably, the Ni/Ca exhibits a stable H₂:CO ratio but a decreased

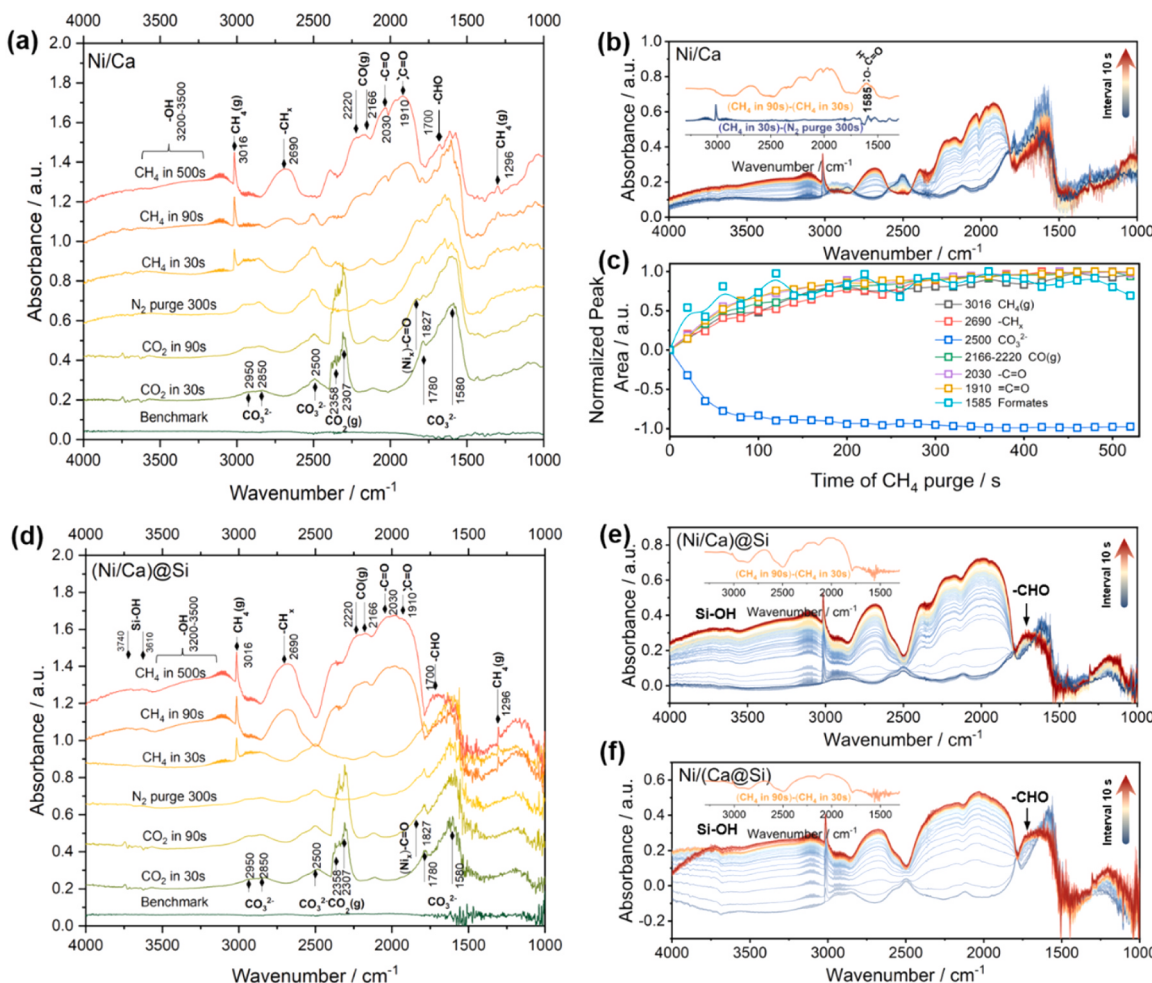


Fig. 9. Isothermal in situ DRIFTS of Ni/Ca material: detailed stacked spectra of ICCU-DRM (a); merged spectra of the integrated DRM step (b); evolvement of normalized selected peak area with time curves (c). Isothermal in situ DRIFTS of (Ni/Ca)@Si and Ni/(Ca@Si) nanoreactors: detailed stacked spectra of ICCU-DRM using (Ni/Ca)@Si nanoreactor (d); merged spectra of the integrated DRM step using (Ni/Ca)@Si (e) and Ni/(Ca@Si) (f) nanoreactors.

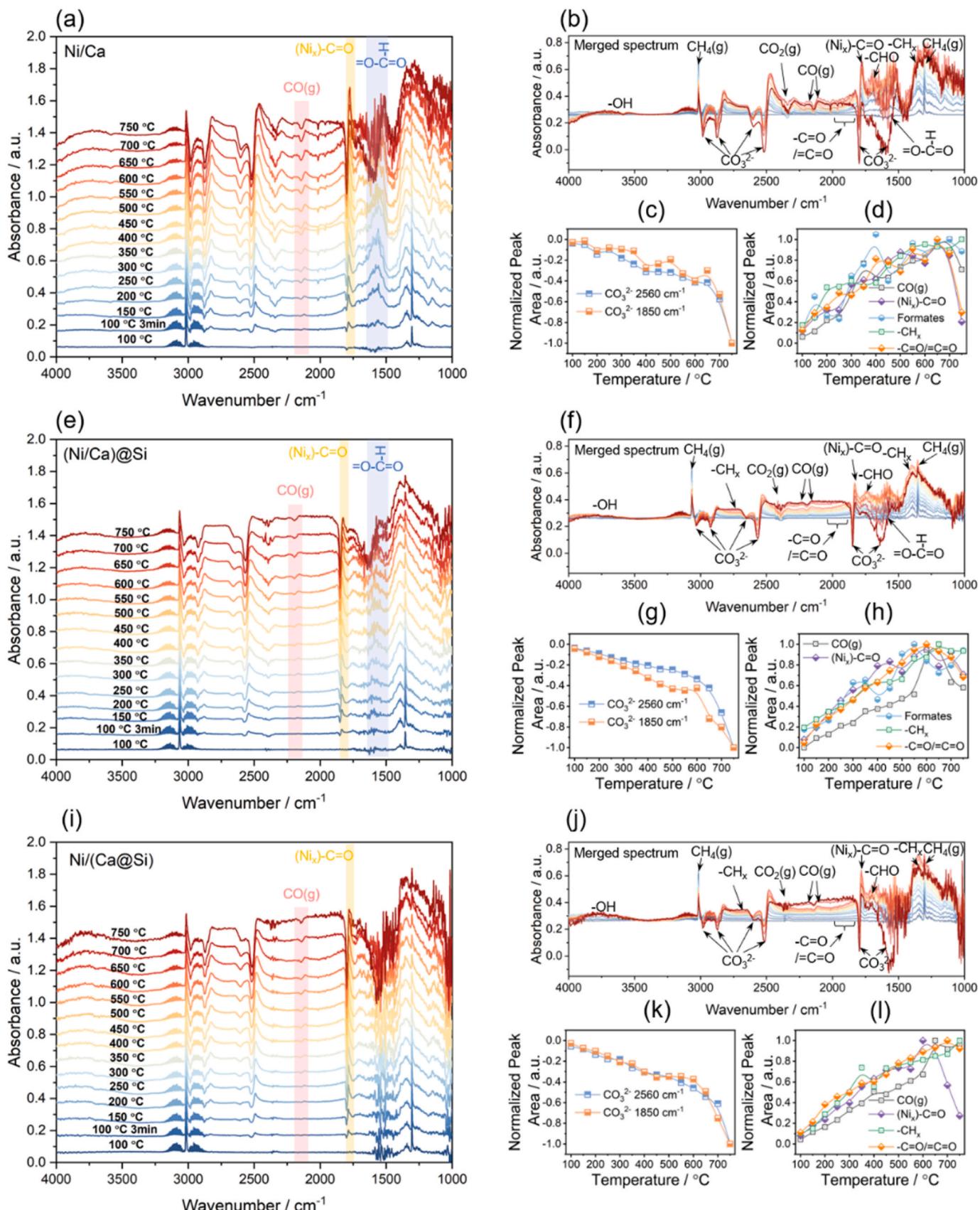


Fig. 10. Stacked and merged DRIFTS spectra of temperature-programmed DRM of (a and b) Ni/Ca, (e and f) (Ni/Ca)@Si and (i and j) Ni/(Ca@Si) nanoreactors. Evolvement of normalized selected peak area with temperature curves of (c and d) Ni/Ca, (g and h) (Ni/Ca)@Si and (k and l) Ni/(Ca@Si) nanoreactors.

carbon deposition amount, which is mainly attributed to the decreased DRM time consumption due to the declined CO₂ capacities in cycles. It was reported that the CaO went through intense dynamic volume change in cyclic carbonation-decomposition[59], which induces the formation of encapsulated Ni. Furthermore, the Ca₂SiO₄ shell also dynamically changes, as evidenced by Fig. 7. In summary, the encapsulation of Ni by Ca₂SiO₄ and CaO is responsible for the gradually decreasing amount of carbon deposition and H₂:CO ratio.

The CO₂ conversion is the most pivotal indicator to evaluate the catalytic performance in the integrated DRM process. It is noted that the (Ni/Ca)@Si nanoreactor outperforms the other two counterparts, as detailed in Fig. 6c. Such results successfully demonstrated the effectiveness of the confined catalysis strategy. For Ni/Ca material, the CO₂ generated from carbonate decomposition most likely misses nearby catalytic sites. For Ni/(Ca@Si), although the decomposed CO₂ was well confined by the Ca₂SiO₄ shell, the residual CO₂ in the cavities had no chance to contact catalysts. Constructing active sites inside the shell would convert the confined CO₂ more thoroughly and hence exhibit enhanced catalytic performance. On the other hand, all three candidates exhibited impressive stabilities with no distinct CO₂ conversion rate decrease. However, as cyclic real-time results shown in Figure S4, the (Ni/Ca)@Si and Ni/(Ca@Si) perform a decrease in instantaneous CO yield, which is highly attributed to the hinderance of the Ca₂SiO₄ shell. It is suggested that the shell structure (e.g. thickness, porosity) should be carefully designed to achieve confined catalysis and satisfactory kinetics.

Consistent with the stable catalytic performance, there is no distinct change in the phase composition (Fig. 8a). Furthermore, the Ca₂SiO₄ phase on Ni/(Ca@Si) exhibits higher crystallinity compared to (Ni/Ca)@Si, as illustrated in the inserted patterns in Fig. 8a. The crystallinity of the Ca₂SiO₄ shell could be clearly evidenced by TEM images in Fig. 7, in which the (Ni/Ca)@Si performs much thinner crystals.

Furthermore, the CaO yolks and Ca₂SiO₄ shells went through dynamic changes in cycles of ICCU, which could be further evidenced by porosity in Fig. 8b. In general, the (Ni/Ca)@Si and Ni/(Ca@Si) nanoreactors perform higher surface areas (Table S2) and more abundant micro/mesopores, which are attributed to the yolk-shell structure. Compared to the as-synthesized materials (Fig. 2d), all the NiSiCa-based materials show better mesopores, even distinctly sintered (e.g. Ni/Ca, Fig. 7a). In previous studies, it was found that the dynamic volume change of CaO-based materials in carbonation/decomposition would induce abundant cracks during sintering[19,60]. Here, we suggest that the sintering of CaO does not always exhibit pore structure loss and a decrease in specific surface area.

The Raman spectrum could characterize the detailed information of carbon species with different orderliness. As illustrated in Fig. 8c, the D band at ~1357 cm⁻¹ and G band at 1579 cm⁻¹ identify the amorphous/disordered carbon and sp² stretching vibration of graphitic carbon species, respectively[61,62]. The intensity ratio of D and G bands was therefore widely applied to characterize the degree of graphitization[63, 64]. As exhibited in Fig. 8c, the Ni/Ca performs the lowest I_D:I_G ratio (0.76) and highest carbon deposition amount (Fig. 6b), indicating the growth of large amounts of graphitized carbon or nanotubes (also evidenced by Fig. 7a). The (Ni/Ca)@Si and Ni/(Ca@Si) nanoreactors are more intent to induce amorphous carbon, of which the Ni/(Ca@Si) exhibits the highest I_D:I_G ratio (1.20). By jointly analyzing the carbon deposit amounts of all three materials (Fig. 8b), it is clear that the formation of amorphous carbon is negatively correlated to the generation of carbon nanotubes. It has been proven an effective strategy by impeding the migration of carbon species on the metal clusters' surface to inhibit the growth of carbon nanotubes[65]. In line with the above-mentioned discussion, the encapsulation extent of Ni is negatively corrected to the carbon deposit amount. The carbon deposition would be gasified into toxic CO in the CO₂ capture step and harm the exhaust gas. Therefore, the best scenario in the integrated DRM step is to selectively catalyze DRM with no CH₄ decomposition. The constructed

nanoreactors deliver such expectations brilliantly with even enhanced catalytic activities (Fig. 6c).

3.4. In situ diffused reflectance infrared fourier transform spectroscopy (DRIFTS)

The reaction mechanism and potential intermediates were studied with in situ DRIFTS under isothermal conditions (650 °C), as shown in Fig. 9. The CO₂ capture step of all the NiSiCa-based materials performs similar species changes. As detailed in Figs. 9a and d, a group of peaks belonging to carbonates emerged and quickly saturated (i.e. 30 s) on the surface during CO₂ capture, including ~ 2950, 2850, 2500, 1780 and 1580 cm⁻¹[19]. The surface reaction of CaO and CO₂ is reported as a chemical reaction-controlled process with fast kinetics[66], explaining the rapid saturation of surface carbonates. To supplement carbonates in the bulk phase for the following DRM, we performed the carbonation step for 90 seconds. Interestingly, as shown in Fig. 9a, a peak at ~ 1827 cm⁻¹ gradually emerged on Ni/Ca after 90 s' CO₂ capture, which is contributed by the formation of multidentate carbonyls (Nix-C=O) [67]. However, the counterpart nanoreactors (Fig. 9d and S6) did not generate such species under the same CO₂ exposure conditions. It was reported that the Ni would decompose CO₂ and be oxidized via carbonyl intermediates[68,69]. The Ni clusters on Ni/Ca are larger and, hence, might be more prone to yield stable multidentate carbonyl species.

The surface carbonates species showed no distinct change after ~ 300 s N₂ purge, nor did stable bridging carbonyl species. Subsequently, the CH₄ was introduced to investigate the integrated DRM process. As shown in Fig. 9a, the peaks at 3016 and 1296 cm⁻¹ are attributed to the C-H stretching and bending vibrations of CH₄, respectively[70]. The integrated DRM is slower compared to surface carbonation in in situ DRIFTS. As evidenced in the inserted patterns in Fig. 9b, no apparent new species formed on the subtracted spectrum after 30 s' CH₄ exposure. Subsequently, several key intermediates emerged with time on Ni/Ca accompanied by surface carbonates consumption (Fig. 9a, b and c). Notably, the consuming carbonates species (~1580 cm⁻¹) and generating formates (~ 1585 cm⁻¹)[70,71] are highly overlapped, and the emerged formates could be identified by deducting the previous spectrum (the insert pattern in Fig. 9b). Moreover, the carbon, formed from CH₄ decomposition, might be oxidized by surface hydroxyl to form -CHO species (~ 1700 cm⁻¹)[72]. Specifically, CHx (2690 cm⁻¹), linearly coordinated carbonyls (-C=O, 2030 cm⁻¹), bridging carbonyls (=C=O, 1910 cm⁻¹)[73], -CHO (1700 cm⁻¹), formates (HCOO, 1585 cm⁻¹) and gaseous CO (2220 and 2166 cm⁻¹) gradually increased and reached the maximum level at ~ 200 s. Subsequently, the bulk carbonates replenish CO₂ to the surface for DRM. Due to the sensitivity of surface characterization, it is difficult to observe the complete disappearance of key species within a limited time (i.e., 500 s). Similar to Ni/Ca, the (Ni/Ca)@Si and Ni/(Ca@Si) nanoreactors exhibit the increasing of carbonyls and CHO species while yielding gaseous CO (Fig. 9d and S6). Differently, the formates species are invisible, which might be masked by the consumption of carbonates (~ 1580 cm⁻¹). It at least proves that the formates formation on nanoreactors is not as significant as Ni/Ca. Furthermore, the (Ni/Ca)@Si (Fig. 9e) and Ni/(Ca@Si) (Fig. 9f) show more distinct CHO groups compared to Ni/Ca (Fig. 9b), which are attributed to the abundant Si-OH groups (~ 3700 cm⁻¹)[74].

The in situ DRIFTS was also conducted at various temperatures to identify the temperature sensitivity of the formation of intermediates. Before temperature-programmed DRIFTS evaluation, the materials were thoroughly carbonated at 650 °C for 30 min. As shown in Fig. 10a and b, the intermediates, including carbonyls, formates and CHO species, could form on Ni/Ca at low temperatures (e.g., 100 °C). The gaseous CO could be generated at ~ 200 °C but with low kinetics. With the temperature increase, the surface carbonates were gradually consumed. The consumption rate of carbonates significantly increases from ~ 650 °C (Fig. 10c, g and k), accompanied by the maximum intermediates peak

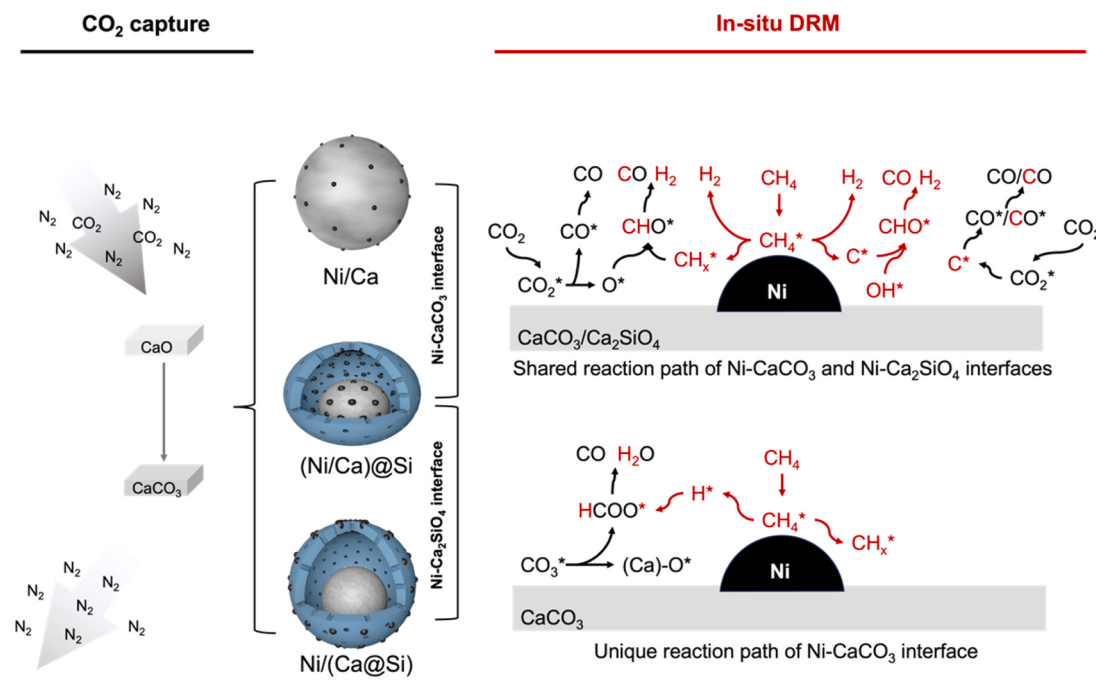


Fig. 11. Mechanism diagram of ICCU-DRM over Ni/Ca, (Ni/Ca)@Si and Ni/(Ca@Si) nanoreactors.

area and CO yields (Fig. 10d, h and i). Importantly, the temperature-programmed DRIFTS identified the various intermediate species on the three candidates. Specifically, carbonyls, formates and CHO species were characterized on Ni/Ca and (Ni/Ca)@Si, while no formates species were detected on Ni/(Ca@Si).

As demonstrated in Fig. 11, the universal mechanism of CO₂ capture is the formation of CaCO₃. Subsequently, the CH₄ decomposes into CH_x/C and H on Ni and releases H₂ by combining two H. The CO₂/CO₃²⁻ could dissociate on Ni surface and yield carbonyl species and then desorbed into gas, simultaneously supplying O for C gasification. The C might also be oxidized by surface OH as an intermediate (CHO) for CO generation. Differently, the Ni/Ca and (Ni/Ca)@Si, with Ni-CaCO₃ interfaces, exhibited the formates intermediates, which are absent on Ni/(Ca@Si). The formate intermediate most likely generates from the direct reaction of spillovered hydrogen with carbonates. In summary, the Ni catalyzes the DRM potentially via carbonyls and CHO intermediates, while the formate intermediate is dependent on the Ni-carbonates interface in the integrated DRM.

4. Conclusion

Inspired by confined catalysis, Ni-functionalized Ca@Si yolk-shell nanoreactors were constructed using novel dry-ice precipitation and silane hydrolysis methods. The Ca₂SiO₄ shell hindered the sintering of CaO and tailored the diffusion of decomposed CO₂ in the integrated DRM process. By doping Ni inside the Ca₂SiO₄ shell, the confined CO₂ could thoroughly react with CH₄ and stably exhibit ~ 92% CO₂ conversion in 10 cycles at 650 °C. The local structure of active sites could be critically tuned by doping Ni in/out of the Ca₂SiO₄ shell to evaluate the catalytic roles of Ni-carbonates or Ni-Ca₂SiO₄ interfaces. Based on in situ DRIFTS investigations, it is found that the integrated DRM could be catalyzed via carbonyls, formates and CHO intermediate steps, while the formates are highly related to the Ni-carbonates interface. The reported confined catalysis directed nanoreactor construction and identified potential intermediates provide valuable reference and portable strategy for further deployments on ICCU and other heterogeneous catalytic processes.

CRediT authorship contribution statement

Zhu Yuan: Investigation. **Sun Shuzhuang:** Writing – review & editing, Writing – original draft, Methodology, Investigation, Formal analysis, Data curation, Conceptualization. **Xu Yongqing:** Writing – review & editing, Supervision, Methodology, Investigation, Formal analysis. **Wu Mengna:** Formal analysis. **Zhang Chen:** Investigation. **Wang Yuanyuan:** Methodology, Investigation, Formal analysis. **Zhou Hui:** Writing – review & editing, Supervision, Resources, Funding acquisition. **Xu Yikai:** Methodology, Formal analysis, Conceptualization. **Wu Chunfei:** Writing – review & editing, Supervision, Resources, Project administration, Funding acquisition, Conceptualization. **Sun Hongman:** Writing – review & editing, Investigation. **Yang Xiaoxiao:** Formal analysis. **Bie Xuan:** Investigation, Formal analysis. **Zhao Xiaotong:** Investigation, Formal analysis. **Zhang Yingrui:** Investigation, Formal analysis.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Data availability

Data will be made available on request.

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Appendix A. Supporting information

Supplementary data associated with this article can be found in the

online version at doi:10.1016/j.apcatb.2024.123838.

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